(22) Filed 12 Aug. 1977 Application No. 33898/77 Convention Application No. 2636425 (32) Filed 13 Aug. 1976 in

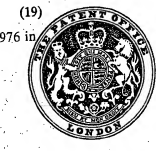
Fed. Rep. of Germany (DE)

H660

Complete Specification Published 28 Jan. 1981

C08F 226/02 246/00 INT. CL.3

Index at Acceptance (52)C2C 200 20Y 30Y 340 34Y 577 627 AA C3Y B230 B240 B241 B245 F530 F550



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## (54) CURABLE COATING COMPOSITIONS COMPRISING VINYLURETHANES

(71) We, BASF AKTIENGESELLSCHAFT, a German Joint Stock Company of 6700 Ludwigshafen, Federal Republic of Germany, do hereby declare the invention, for which we pray that a Patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following Statement:-

The present invention relates to a curable coating composition which comprises a vin-

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Coating compositions comprising solutions of high molecular weight or low molecular ylurethane. weight unsaturated compounds in vinyl monomers have been disclosed. If vinyl compounds which copolymerize rapidly, e.g. acrylic compounds, are used, such systems can be cured with high energy radiation. However, acrylic compounds have the disadvantage that they are

usually toxic and cause severe irritation of the skin. A process has been disclosed in German Laid-Open Application DOS 2,064,701 for the manufacture of coatings by curing olefinically unsaturated polymeric materials containing urethane groups, these materials, with or without admixture of other olefinically unsaturated monomeric compounds, being cured by means of ionizing rays; the olefinically unsaturated polymeric material containing urethane groups comprises at least two CH<sub>2</sub>=CH-NH-C-O-

units and has been obtained by reacting hydrolic oligomers or polymers of molecular weight from 500 to 10,000 with vinyl isocyanate. However, these products are rather unsuitable for certain applications, where a particularly high degree of crosslinking is required. In some cases, a very low viscosity of the component which forms the binder is desirable, to facilitate

The present invention therefore seeks to provide improved coating compositions particularly those which, in spite of being processed at a low viscosity, cure very actively. We have found that good results can be achieved by using special mixtures of certain vinyl

urethanes and other unsaturated compounds. The present invention provides a curable coating composition which comprises a mixture of A) one or more olefinically unsaturated compounds having a molecular weight of from 70

to 20,000 and a boiling point at standard pressure of above 50°C and B) one or more vinylurethanes copolymerizable with A) but differing therefrom, in which component (B) is a reaction product of vinyl isocyanate and a polyol having a molecular weight of less than 500 and the weight ratio of the components A:B is from 99:1 to

The following details relating to the components on which the coating compositions of the 30:70.

(A) The olefinically unsaturated compounds (A) should have a boiling point at standard pressure of above 50°C, preferably above 100°C. Lower-boiling compounds are, in general, less suitable, since they evaporate too easily after applying the coating compositions and before curing them. This can alter the analysis of the coating composition and can also lead to pollution of the environment by malodorous and, in some cases, toxic monomers. Compounds having a vapor pressure of less than 10 mm Hg at 100°C are therefore particularly preferred for finishes and printing inks which do not pollute the environment.

The molecular weight of component A can vary within wide limits, from 70 to 20,000, i.e. either low molecular weight or high molecular weight compounds may be employed. The compounds may be mono-olefinically unsaturated or have a higher degree of unsaturation.

		Examples of suitable components A are:  1. Esters of unsaturated monocarboxylic acids or dicarboxylic acids, for example esters of acrylic acid, methacrylic acid, $\alpha$ -cyanoacrylic acid, crotonic acid, cinnamic acid, sorbic acid, maleic acid, fumaric acid or itaconic acid, with aliphatic, cycloaliphatic or aromatic-aliphatic	
	5	monohydric to tetrahydric alcohols of 3 to 20 carbon atoms, e.g. methyl acrylate and methacrylate, n-, i- and t-butyl acrylate and methacrylate, 2-ethylhexyl acrylate, lauryl acrylate, dihydrodicyclopentadienyl acrylate and methacrylate, methylglycol acrylate, hydroxypropyl acrylate and methacrylate, ethylene	<b>5</b> .
	10	glycol diacrylate, diethylene glycol diacrylate, triethylene glycol diacrylate, neopentyl-glycol diacrylate and dimethacrylate, 1,4-dimethylolcyclohexane diacrylate, pentaerythritol triacrylate, tetraacrylate, trimethacrylate and tetramethacrylate, ethyl $\alpha$ -cyanoacrylate, ethyl crotonate, ethyl sorbate, diethylmaleate, diethyl fumarate and the diacrylate and dimethacrylate of oxyalkylated bisphenol A.	10
	15	2. Amides of acrylic acid or methacrylic acid which may or may not be substituted by alkyl, alkoxyalkyl or hydroxyalkyl at the nitrogen, e.g. N,N'-dimethylacrylamide, N-isobutylacrylamide, diacetoneacrylamide, N-methylolacrylamide, N-methylacrylamide, N-butoxymethylacrylamide, N-butoxymethylacrylamide	15
	20	and ethylene glycol bis-(N-methylolacrylamide) ether.  3. Vinyl esters of monocarboxylic acids or dicarboxylic acids of 2 to 20 carbon atoms, e.g. vinyl acetate, vinyl propionate, vinyl 2-ethylhexanoate, vinyl versatate ("Versatic" is a positioned Trade Mark) or divinyl adjuste.	20
,	25	4. Vinyl ethers of monohydric or dihydric alcohols of 3 to 20 carbon atoms, e.g. isobutyl vinyl ether, hexyl vinyl ether, octadecyl vinyl ether, ethylene glycol divinyl ether, diethylene glycol divinyl ether, butanediol divinyl ether and hexanediol divinyl ether.  New New York and Divinyl compounds as a New invloyer collidore. New invloying income.	25
		N-vinylcaprolactam, N-vinylmorpholine, N-vinyloxazolidone, N-vinylsuccinimide, N-methyl-N-vinylformamide, N-vinylcarbazole and divinylureas, e.g. N,N' δ divinylethyleneurea and divinylpropyleneurea.  6. Styrene and its derivatives, e.g. α-methylstyrene, 4-chlorostyrene and 1,4-	30
	30	divinylbenzene.  7. Allyl ethers and allyl esters, e.g. trimethylolpropane diallyl ether, trimethylolpropane triallyl ether, pentaerythritol triallyl ether, diallyl maleate, diallyl fumarate and diallyl phthalate.	
	35	8. Unsaturated polyesters having a molecular weight of from 500 to 5,000 and containing from 0.5 to 10 double bonds per 1,000 molecular weight units, the polyesters being manufactured from, for example, a) from 10 to 70 per cent by weight of one or more α.β-unsaturated dicarboxylic acids, e.g.	35
	40	malcic acid, fumaric acid or itaconic acid, b) from 0 to 60 per cent by weight of one or more saturated aliphatic, optionally chlorine- substituted cycloaliphatic or aromatic dicarboxylic acids, e.g. succinic acid, adipic acid, tetrahydrophthalic acid, hexahydrophthalic acid, hexachloroendomethylenetetrahydro- phthalic acid, cyclohexane-1, 4-dicarboxylic acid, phthalic acid, isophthalic acid or tereph-	40
	45	thalic acid, c) from 20 to 80 per cent by weight of one or more aliphatic, cycloaliphatic or non-phenolic aromatic diols, e.g. ethylene glycol, diethylene glycol, triethylene glycol, propane-1,2-diol, propane-1,3-diol, butane-1,2-diol, butane-1,3-diol, butane-1,4-diol, neopentyl-glycol, hexane-1,6-diol or oxyalkylated bisphenol A,	45
	50	d) from 0 to 5 per cent by weight of one or more tricarboxylic acids or tetracarboxylic acids, e.g. trimellitic acid, pyromellitic acid or benzenetetracarboxylic acid, e) from 0 to 10 per cent by weight of one or more monocarboxylic acids, e.g. acetic acid, propionic acid or benzoic acid, f) from 0 to 5 per cent by weight of one or more trifunctional or tetrafunctional alcohols, e.g.	50
	55	glycerol, trimethylolpropane or pentaerythritol, and g) from 0 to 10 per cent by weight of one or more monofunctional alcohols, e.g. methanol, ethanol, propanol or butanol.  9. Unsaturated epoxy resins which have been manufactured from, for example, a) a monofunctional epoxide and acrylic acid or methacrylic acid, as described in U.S.	55
	60	Patent 2,484,487. b) a bifunctional epoxide and an unsaturated fatty acid, as described in U.S. Patent 2,456,408. c) a polyfunctional aromatic epoxide and crotonic acid, as described in U.S. Patent 2,575,440 or	: <b>6</b> 0
	65	d) a polyfunctional aromatic or aliphatic glycidyl ether and acrylic acid or methacrylic acid, as described in U.S. Patent 2.824.851.  10. Unsaturated polyurethanes obtained from hydroxyalkyl acrylates and disocyanates.	· 65
	65	10. Chattilated polydrechanies dotained from hy	

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	with or without polyols or polyamines, for example saturated or unsaturated polyester- with or without polyols or polyamines, for example saturated or unsaturated polyester- with or without polyols or polyamines, for example saturated or unsaturated polyester- with or without polyols or polyamines, for example saturated or unsaturated polyester-		
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	Laid-Open Application DOS 7, 358 048	5	
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5	11. Unsaturated copolymers, for example manufactured by leaching.  a) copolymers, containing maleic anhydride groups, with unsaturated alcohols, for example  a) copolymers, Containing maleic anhydride groups, with unsaturated alcohols, for example  a) copolymers, Containing maleic anhydride groups, with unsaturated alcohols, for example		
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	as described in German Laid-Open Application DOS 2,050,895 of as described in German Laid-Open Application DOS 2,050,895 of		
	h) acrylic ester copolymers of polyesters, contament		
	unsaturated epositions, e.g. glycidyl acrylate. unsaturated epositions, e.g. glycidyl acrylate.	10	
10	unsaturated epoxides, e.g. glycidyl acrylate.  12. Butadiene polymers in which the double bonds are predominantly present as vinyl		
-	side groups.		
	12 Digital phthalate preparations.		
•	Of course, mixtures of the various unsaturated compounds A may also be used.  Of course, mixtures of the various unsaturated compounds I may also be used.  The compounds listed under 1, 2, 8, 9, 10 and 11 above, and their mixtures, are particularly		
	The compositors listed under 1,2,0,3,2	15	
15	mediatred continuousling A		
IJ.	(B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (B) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component (B) are reaction processes.  (C) The vinylurethanes which may be used as component		
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	than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of suitable polyols are linear or branched animatic, cyclodar than 400. Examples of		
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20	mothyl 2 propyl-propane-1,3-diol, decanediol, cyclonolical diols containing ether groups.		
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	a distribution divicit. Illiculty out 5-1		
	arido and atabulene unide and addition and additional and additional and all and a second and aridonal and additional additional and additional addition	25	
0.5	dihydric phenols; triols, e.g. glycerol, trimethylolpropane, bittane-1,2,4-triol, particle of tribydric alcohols of phenols; alcohols of higher functionality, e.g.	-	
25	1,2,5-triol, hexane-1,2,6-triol, tris-hydroxyethyl isocyanurate, and oxyethylation 1,2,5-triol, hexane-1,2,6-triol, tris-hydroxyethyl isocyanurate, and oxyethylation allowed trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcohols of higher functionality, e.g. ropylation products of trihydric alcohols or phenols; alcoh		
	1,2,5-thirth house sof trihydric alcohols or phenols; alcohols of light intercently, 100 and overroughtion		
. , .	pentaerythritol, dipentaerythritol, sorbitol, mannitol and oxyethylation and oxyethy	30	
	ypivalic acid ester of neopentyl-glycol, and condensation products of dicarooxylic acids or excess diol, which may or may not contain co-condensed higher-functional carboxylic acids or excess diol, which may or may not contain co-condensed higher-functional carboxylic acids or		
30			
	-1bolo		
	alcohols.  To manufacture the vinylurethanes the polyol(s) are reacted with vinyl isocyanate, in general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such an amount as to provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such as the provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such as the provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such as the provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such as the provide from 0.9 to 1.05, moles of general in such as the provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such as the provide from 0.7 to 1.2, preferably from 0.9 to 1.05, moles of general in such as the provide from 0.9 to 1.05, moles of general in such as the provide from 0.9 to 1.05, moles of general in such as the provide from 0.9 to 1.05, moles of general in such as the provide from 0.9 to 1.05, moles of general in such as the provide from 0.9 to 1.05, moles of general in such as the provide		
	and in such an amount as to provide from 0.7 to 1.2, preferably from 0.7 to 1.05, more	35	
<u>ــ</u>	general in such an amount as to provide from 0.7 to 1.2, per mole of OH groups. vinyl isocyanate per equivalent of polyol, i.e. per mole of OH groups.		
35	TARILLANDAR AT TRACTION DI ULIGIS OF TANDES OF		
	used as component (B).  The properties of the products can, to a certain degree, be suited to special requirements by  The properties of the starting components.		
	The properties of the products can, to a certain degree, be suited to special requirements		
	The properties of the products day, appropriate variation of the starting components.  appropriate variation of the starting components.	40	
	Composite A sin B ale employed in the		•
40	from 95:5 to 50:50.  In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, components (A) and (B) are mixed at from 20 to 80°C in conventional mixing In general, conventional stirring equipment.		
	In general components (A) and (B) are mixed at from 20 to 80°C in convenients		
	In general, components (A) and (S) and (S) are equipment, or using conventional stirring equipment, or using conventional stirring equipment.  The coating composition of the invention may, depending on its intended use, also contain the coating composition of the invention may.		
•		45	
45	1. From 0 to 70, preferably from 10 to 50, per cent by weight of inorganic of inorganic of the pigment, e.g. carbon black, titanium dioxide, chalk, baryte, zinc white, lithopone, chromium pigment, e.g. carbon black, titanium dioxide, chalk, baryte, zinc white, lithopone, chromium pigment, e.g. phthalocyanines, azo pigments, anthra-		
	rigment e g carbon black, titanium dioxide, chalk, baryte, zhe white, interpolation and pigments, anthra-		
	TO BOTH THE HALL OF COMPLETE P.D.		
	quinone colorants and quinacridone pigments.  Quinone colorants and quinacridone pigments.  2. From 0 to 10, preferably from 0.001 to 3, per cent by weight of dye, e.g. eosin, crystal	50	
	a 4 Erom II to III Dreiciauly montour		
51	violet or malachite green.  3. From 0 to 10, preferably from 1 to 5, per cent by weight of leveling agent, e.g. butyl		
	2. From 0 to 10 preferably from 1 to 5, per cent by weight of leveling agont of		
	acetate, butanol, silicones or thinners.		
	acetate, butanol, silicones or thinners.  4. From 0 to 1, preferably from 10 to 50, per cent by weight of inhibitor.	55·	
	4. From 0 to 1, preferably from 0.001 to 0.3, per cent by weight of filler.  5. From 0 to 70, preferably from 1 to 50, per cent by weight of diluent.	••,	
5	5. From 0 to 70, preferably from 10 to 50, per cent by weight of diluent. 6. From 0 to 10, preferably from 10 to 50, per cent by weight of inert synthetic resin, e.g.	•	
	6. From 0 to 10, preferably from 1 to 5, per cent by weight of undert. 7. From 0 to 70, preferably from 10 to 50, per cent by weight of inert synthetic resin, e.g.		
	aminoplasts and alkyd resins.		
	aminoplasts and alkyd resins.  8. From 0 to 7, preferably from 0.5, to 5, per cent by weight of wax.  8. From 0 to 7, preferably from 0.2 to 5, per cent by weight of thixotropic agent.	60	
	8. From 0 to 7, preferably from 0.5, to 5, per cent by weight of thixotropic agent. 9. From 0 to 5, preferably from 0.2 to 5, per cent by weight of thixotropic agent.	,	
6	9. From 0 to 5, preferably from 0.2 to 5, per cent by weight of thixotropic age. This may The coating compositions are cured by copolymerization of components A and B. This may The coating compositions are cured by means of infrared radiation or with the aid of ar	1	
	The coating compositions are cured by copolymerization of compositions are actived by means of infrared radiation or with the aid of are be effected thermally, by direct heating, by means of infrared radiation or with the aid of are being the professibly by using high energy radiation.		
	be effected thermally, by diffect hearing, or initiator, but preferably by using high energy radiation. initiator, but preferably by using high energy radiation.	-	
	initiator, but preferably by using high energy radiation.  For the thermal polymerization, temperatures of from 20 to 200°C are generally emp.  For the thermal polymerization, temperatures or azo compounds, and accelerators	65	
		. 05,	
6	55 loyed; conventional initiations, org.		

5	e.g. cobalt salts, may be added in amounts of from 0.5 to 6 per cent by weight.  Radiation curing may be effected with UV rays or by electron beams. In the former case, a conventional photoinitiator, such as are described, for example, by B.J. Kosar in "Light Sensitive Systems", Wiley, 1965, pages 158-193, may be added. Preferably, the following are used: Benzoin, benzoin ethers, diacetyl, benzil, benzil monoketals, benzophenone, Michler's ketone, xanthones, anthraquinones, sulfur compounds, e.g. disulfides, thiols and dithiocar-	5
l <b>0</b> ်	bamates, and carbonyl compounds, e.g. triphenylphosphine-iron tetracarbonyl in conjunction with chlorine donors, each in amounts of from 1 to 3 per cent by weight. Details of the method of curing by UV radiation are to be found in "Photopolymerization" by H. Barzynski, K. Penzien and O. Volkert in Chemiker-Zeitung 96 (1972), 545-551, and in German Laid-Open Application DOS 2,251,933. Electron beam curing is described in detail in German Laid-Open Application DOS 2,049,715 cited above, in which further details regarding advantageous coating thicknesses, radiation doses and irradiation times may also be	10
15	found.  The coating compositions may be used for the production of coatings and finishes on metals, wood, plastic, leather and paper. These coatings may be used as protective layers or	15
20	for decorative purposes.  The coating compositions may also be used as a photopolymer layer for print carriers, e.g. relief print plates or planographic print plates, or for photoresists. In such cases, they are cured by image-wise irradiation of the coated carrier; no curing occurs in the unexposed representation of the layer are dissolved out again.	20
25	Finally, the coating compositions may also be used for the manufacture of UV-curing printing inks and print pastes. Using such inks and pastes, substrates, e.g. paper, metals or plastic films, are printed with the pigment-containing binders in order to apply a layer from about $0.5$ to $5 \mu m$ thick. The binders cure very rapidly on ultraviolet irradiation and the ink becomes fixed to the substrate. The printing inks manufactured using the coating compositions of the invention are distinguished by extremely rapid drying even if only 1 or 2 ultraviolet lamps are used in the case of four-color printing, and by very high printons is particular.	25
30	The drying prints have very good scuff resistance. This would seem to be more in line with what is needed in multi-color printing. This resistance is much improved over conventional ultraviolet printing inks in the case of colors which are, from this point of view, particularly critical in ultraviolet printing, e.g. blue and black. These remarks apply particularly to prints are dueed at maximum print speeds and hence with minimum irradiation times.	30
35	If no pigments are added, the binders may be used as overprinting varnishes or as clear coatings.  In the Examples, parts and percentages are by weight.	35
	EXAMPLE 1	
40	0.07 part of dibutyl-tin dilaurate is added to a solution of 134 parts of trimethylolpropane in 300 parts of ethyl acetate and 207 parts of vinyl isocyanate are added dropwise in the	40
45	course of 40 minutes at from 50 to 60°C, whilst stirring. The fraction is their another to continue for from 1 to 2 hours, until the isocoyanate content has fallen to below 0.1%. The mixture is then cooled to about 10°C, whereupon the reaction product substantially cystallizes out, and the crystals are filtered off. After recrystallization from ethyl acetate and drying under reduced pressure at room temperature. 320 parts of trimethylolpropane-trivinylurethanc are obtained. A mixture of 50 parts of this material and 50 parts of	45
50 ·	board and is cured with 320 KV electrons at a belt speed of 100 m/mm, corresponding to a	50
	EXAMPLE 2	
<b>5</b> 5	If the procedure of Example 1 is followed, but instead of 134 parts of trimethylolpropane 204 parts of the neopentylglycol ester of hydroxypivalic acid are used, and instead of 207 parts only 138 parts of vinyl isocyanate are used to manufacture component (B), a viscous mass, which solidifes after from 1 to 2 days, is obtained after removing the solvent. A mixture	55
60	of 20 parts of this reaction product with 40 parts of butane-1,4-diol diacrylate and 40 parts of unsaturated polyester obtained from 2 moles of maleic anhydride, 1 mole of hexachloroendomethylenetetrahydrophthalic acid, 2 moles of neopentyl-glycol and 1 mole of propylene glycol (acid number: 30 mg of KOH/g) is applied, as described in Example 1, as a layer 80 um thick to a surface-filled wooden board, and cured with 320 KV electrons at a	60
65	belt speed of 60 m/min, corresponding to a dose of 1.17 Mrad.  A very hard, scratch-resistant coating (König pendulum hardness: 205 seconds) is obtained.	65

## EXAMPLE 3

	Charton 1.4	
	A mixture of 10 parts of trimethylolpropane-trivinylurethane, 40 parts of butane-1,4- oldiacrylate and 50 parts of a reaction product of one mole of a diglycidyl ether, based on oldiacrylate and 50 parts of a reaction product of one moles of acrylic acid is applied, as	5
	A mixture of 10 parts of a reaction product of one mole of a digryclay editor, oldiacrylate and 50 parts of a reaction product of one mole of a digryclay editor, as sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is applied, as sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is applied, as sphenol A (epoxide equivalent weight: 340) with 2 moles of a digryclay editor, as	
5	sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately sphenol A (epoxide equivalent weight: 340) with 2 moles of acrylic acid is approximately acid in a constant weight: 340) with 2 moles of acrylic acid is approximately acid in a constant weight: 340 per acid in acid in a constant weight: 340 per acid in acid	
	escribed in Example 1, as a layer 10 $\mu$ m thick to a saturated, tack-free coating has a Komg	
	sphenol A (epoxide equivalent) $\mu_m$ thick to a surface-filled find fiberootis as a König escribed in Example 1, as a layer 70 $\mu_m$ thick to a surface-filled find fiberootis as a König of the conditions described in Example 2. The cured, tack-free coating has a König of the conditions described in Example 2. The cured, tack-free coating has a König of the conditions of 200 seconds.	0
, '	endulum hardness of patent application No. 33899/// (Serial No. 1363 No.	· .
10	Our copending GB patent application 100.  Our copending GB patent application 100.  aims curable binders for coating compositions comprising urethane groups, which are a laims curable binders for coating compounds containing urethane groups, which are a laims curable binders for coating at least one hydroxyl	
	Our copending GB patched appropriate appropriate accompositions comprising laims curable binders for coating compositions containing urethane groups, which are a lone or more olefinically unsaturated compounds which contains at least one hydroxyl one or more object of vinyl isocyanate and a compound which contains at least one hydroxyl one of vinyl isocyanate and a compound which contains at least one hydroxyl one of vinyl isocyanate and a compound which contains at least one hydroxyl or contains at least one h	:
	laims curable bliders to could be a compound containing uretnate groups, which one or more olefinically unsaturated compound which contains at least one hydroxyl eaction product of vinyl isocyanate and a compound which contains at least one hydroxyl eaction product one radical of an ester of an $\alpha$ , B-olefinically unsaturated carboxylic acid of	
	one or more definitionly unitarity and a compound which contains at least one by the eaction product of vinyl isocyanate and a compound which contains at least one by the eaction product of vinyl isocyanate and a compound which contains at least one by the eact of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ , B-olefinically unsaturated carboxylic acid of roup and at least one radical of an estern of an $\alpha$ -olefinity and $\alpha$ -olefinity an	15
15	roup and at least one radical of all esters of all esters of the carbon atoms, and optionally also comprising to 6 carbon atoms, and optionally unsaturated compounds which are copolymerizable with ii) one or more further olefinically unsaturated compounds which are copolymerizable with	
15	ii) one or more that olefinically unsaturated compounds which are copyrimetrizing wether groups.	
	ii) one or more luttief of the olefinically unsaturated compounds which are copositions but different from the olefinically unsaturated compound(s) containing urethane groups. Out different from the olefinically unsaturated compound(s) containing urethane groups. We make no claim herein to coating compositions comprising components (i) and (ii) We make no claim herein to from 70:30 to 1:99.	
	We make no claim herein to coating compositions comprised	20
20	hove in a weight latte of (1). (2)	
20	Subject to this disclaimer,	
	WHAT WE CLAIM IS:-	
<i>:</i> :	WHAT WE CLEARLY 22	25
	1. A curable coating composition which comprises a mixture of  1. A curable coating composition which comprises a mixture of  A) one or more olefinically unsaturated compounds having a molecular weight of from 70 to  A) one or more obling point at standard pressure of above 50°C and  A) but differing therefrom,	
25		
	20,000 and a bonning physicathanes conclumerizable with A) but different and a polyol having	•
	20,000 and a boiling point at standard polymerizable with A) but differing the circle and a polyol having B) one or more vinylurethanes copolymerizable with A) but differing the circle and a polyol having in which mixture component (B) is a reaction product of vinyl isocyanate and a polyol having in which mixture component (B) is a reaction product of the components A:B is from 99:1	30
	B) one or more vinyinternation to the component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which mixture component (B) is a reaction product of vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl isocyanate and a polyonal in which is a vinyl	
30	to 30:70.	
	2. A composition as chained in state of 100°C	•
	of less than 10 initial and allowed in claim 1 or 2, in which component acres by vice acids or	35
	3. A composition as claimed in description as claimed in description as claimed in description as claimed in description as a claimed in descr	
35	dicarboxylic acids with aliphatic, cycloanplatic of acrylic acid or methacrylic acid which	
	rahydric alcohols of authorityted by alkyl, alkoxy-alkyl of hydroxyday, and containing from	
	may or may not be sharing a molecular weight of from 500 to 5,000 energy resing unsatu-	40
	may or may not be substituted unsaturated polyesters having a molecular weight of from 500 to 5,000 and containing unsaturated polyesters having a molecular weight units, unsaturated epoxy resins, unsaturated to 10 double bonds per 1,000 molecular weight units, unsaturated epoxy resins, unsaturated polyurethanes obtained from hydroxyalkyl acrylates and dissocyanates with or without rated polyurethanes obtained from hydroxyalkyl acrylates and dissocyanates with or without rated polyurethanes reaction products of copolymers containing maleic anhydride groups	
40	0.5 to 10 double bonds per 1,000 methods and disocyanates with the rated polyurethanes obtained from hydroxyalkyl acrylates and disocyanates with the rated polyurethanes obtained from hydroxyalkyl acrylates and disocyanates with the rated along the polyurethan acrylates and reaction products of acrylates are restated along and reaction products of acrylates are restated along the polyurethan acrylates.	
.*	polyols or polyalities and reaction products of actylic ester copyrights	
·	with unsaturated above and groups with olefinically unsaturated control which the polyol used to	45
	with unsaturated alcohols, and reasonable with olefinically unsaturated epoxides. containing carboxylic acid groups with olefinically unsaturated epoxides. containing carboxylic acid groups with olefinically unsaturated epoxides. containing carboxylic acid groups with olefinically unsaturated epoxides.  4. A composition as claimed in any of claims 1 to 3, in which the polyol used to 4. A composition as claimed in any of claims 1 to 3, in which the weight ratio of	٠,
4.	4. A composition as claimed in any of claims 1 to 3. In which the weight ratio of manufacture the component (B) has a molecular weight of less than 400.  5. A composition as claimed in any of claims 1 to 4, in which the weight ratio of the second of the	
	5. A composition as claimed in any or standing of the	
•	components A.B is from a laimed in claim 1 and substantially as described in the	50
	foregoing Examples.	-
5	6. A composition as claimed in claim.  foregoing Examples.  7. A composition as claimed in any preceding claim when coated on a surface and cured to the composition as claimed in any preceding claim when coated on a surface and cured to the composition as claimed in any preceding claim when coated on a surface and cured to the cured to the coated on a surface and cured to the cure	
•	by irradiation by incaris of the photographymer layer for a print succession	
	8. A printing ink or printing paste, or a photopolymen 1 to 6. a curable composition as claimed in any one of claims 1 to 6.  J.Y. & G. W. JOHNSON.	55
	a curable composition as a sum I Y & G. W. JOHNSON.	
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	Agents for the Applicants.	60
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